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(54) ANTISENSE BASE SEQUENCES

(57) The present invention provides a novel method to enhance an ability to suppress in vivo protein synthesis in a method of suppressing synthesis by use of an antisense nucleotide sequence. More specifically, the present invention provides an antisense nucleotide sequence comprising two or more successive repeats of the structural gene of interest or a fragment thereof in the antisense direction, an expression vector comprising said antisense nucleotide sequence, a host transformed by said expression vector and a method for suppressing the expression of a protein by use of said antisense nucleotide sequence.

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Description

BACKGROUND OF THE INVENTION

5 [0001] This invention relates to a technique for suppressing protein synthesis using antisense nucleotide sequences, particularly to a technique capable of enhanced suppression of protein synthesis.

PRIOR ART

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[0002] It is known that functional RNAs such as messenger RNA which provides information on protein synthesis are suppressed by RNAs having complementary nucleotide sequences to the aforementioned RNA, which are collectively referred to as antisense RNAs. Research is under way to create plants wherein antisense RNA is introduced artificially by gene recombinant technology. Genes for expressing antisense RNAs are constructed in such a manner that a sequence corresponding to a partial or full length of a DNA sequence (cDNA or genomic DNA) coding for a protein of interest is connected downstream of a promoter in the antisense direction.

[0003] While various antisense techniques have been proposed, the following are worth mentioning:

- (1) Recombinant petunia was prepared that produced an antisense RNA against RNA of a chalcone synthase gene participating in the synthesis of flower pigments, and the recombinant petunia presented a different flower color than the wild type (EP 34-1885A);
- (2) The expression of a polygalacturonase gene, a key factor in tomato fruit losing its firmness, was suppressed by an introduced antisense RNA to create tomatoes that could be preserved for a longer period than the wild type (EP 3891115A):
- (3) Melton et al. used a full-length sequence of β-globin cDNA as an antisense gene (Proc. Natl. Acad. Sci. USA 82:144-148 (1985));
- (4) Stockhaus et al. used as an antisense gene a sequence corresponding to the full length of a 10 kD protein cDNA which takes part in the photo-system of photosynthesis (the EMBO Journal 9:3013-3021 (1990));
- (5) Alexander et al. used as an antisense gene a sequence corresponding to the full length of chalcone synthase cDNA which takes part in the synthesis of flower pigments (Nature 333:866-869(1988));
- (6) Hamilton et al. used a sequence corresponding to the full length of ethylene synthase (ACC-oxidase) cDNA as an antisense gene (Nature 346:284-287 (1990)); and
- (7) Smith et al. used a sequence corresponding to a partial length of polygalacturonase cDNA as an antisense gene (Nature 334:724-726 (1988)).

[0004] Thus, in the conventional antisense-related technology, a nucleotide sequence corresponding to a partial or full length of the nucleotide sequence coding for a protein of interest is simply inserted as an antisense gene in the reverse direction at a downstream of a promoter.

[0005] While several reports have been published which propose methods for reducing the content of a protein of interest using antisense RNA, there is no report of success in reducing protein synthesis when the protein is one of those occurring abundantly in vivo as exemplified by the storage proteins in plant seeds. In order to ensure that the content of a protein occurring abundantly in vivo is reduced using an antisense RNA, the latter must be provided in a large amount at the site of its syntheses. One possible means of satisfying this need is to introduce the antisense gene in multiple copies; however, if homozygote is desired as in the case where the host is a plant, the problem is that a great amount of work is required to fix a gene introduced in multiple copies by self-fertilization. Another possible means is to enhance the activity of a promoter that expresses an antisense gene of interest, but this also is not easy to accomplish.

[0006] The antisense genes reported thus far do not provide an easy way to reduce the content of proteins that occur abundantly in vivo.

SUMMARY OF THE INVENTION

[0007] It is generally believed that the degree of suppression of protein synthesis increases with the increasing expression of associated antisense genes (Melton D.A. et al. (1985) Proc. Natl. Acad. Sci. USA 82, 144-148; and Ecker J.R. et al. (1986) Proc. Natl. Acad. Sci. USA 83, 5372-5376). As already mentioned, the expression of an antisense gene could be increased by, for example, enhancing the activity of a promoter used in expressing the antisense gene or by introducing it in an increased copy number. However, both methods are subject to the problems described above. The purpose of the present invention is to provide an antisense gene that can not only suppress the synthesis of a protein in vivo more efficiently but also lower even the content of a protein that occurs abundantly in vivo.

[0008] If a nucleotide sequence corresponding to a full length of a structural gene of interest is introduced in the anti-

sense direction, the effectiveness in reducing the synthesis of a protein of interest is indeed increased to some extent. However, according to this method in which a nucleotide sequence corresponding to the full length of the structural gene of interest is introduced in the antisense direction, it is possible that an unknown open reading frame will appear, thereby inducing the expression of an unexpected protein in the host. Particularly in the case of edible plants, safety to humans and changes in taste are two important factors that must be considered seriously. The problem of the expression of an unexpected protein can be solved by suppressing it using a sequence corresponding to a portion of the structural gene, namely, a partial sequence which lacks an open reading frame or one from which an open reading frame has been deleted artificially. However, if a part of the structural gene of interest is introduced as a single unit, the effectiveness in reducing the synthesis of the protein by the structural gene will not be attained at the intended level. Under the circumstances, the present inventors conducted intensive studies in order to develop a method for solving the aforementioned problems by reducing the synthesis of a protein of interest without expressing any unknown protein. As a result, the inventors found a method wherein an antisense nucleotide sequence is prepared so that it comprises a plurality of the structural gene of interest or a fragment thereof linked in succession, and said sequence is introduced into a genomic gene in a target cell; the present invention has been accomplished on the basis of this finding.

BRIEF DESCRIPTION OF THE DRAWINGS

[0009]

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- Fig. 1 shows schematically the structures of genes used in the in vitro RNA syntehsis;
 - Fig. 2 shows electrophoretic patterns in the analysis of translation products in a wheat germ extract;
 - Fig. 3 is a graph showing the changes in the amount of glutelin synthesis;
 - Fig. 4 shows schematically the structure of a plasmid vector used in transformation;
 - Fig. 5 shows an electrophoretic pattern in the PCR analysis of transformants having a sequence comprising 8 repeats of glutelin A antisense gene introduced therein;
 - Fig. 6 shows an electrophoretic pattern in the PCR analysis of transformants having a full length of glutelin A antisense gene introduced therein;
 - Fig. 7 shows an electrophoretic pattern in the northern analysis of transformants having a sequence comprising 8 repeats of glutelin A antisense gene introduced therein; and
- 50 Fig. 8 shows schematically the structure of the plasmid vector used in the transformation.

DETAILED DESCRIPTION OF THE INVENTION

[0010] The present inventors have found that an antisense nucleotide sequence may be formed such that it comprises two or more units of a structural gene of interest or a sequence representing a part of said structural gene in succession in the antisense direction, and said sequence is introduced into a genomic gene in a target cell so as to suppress the intracellular expression of a protein encoded by the structural gene effectively. The present invention was accomplished on the basis of this finding.

[0011] Thus, according to one aspect of the invention, there is provided an antisense nucleotide sequence in which two or more units of a structural gene of interest or a sequence representing a part of said structural gene are linked in succession in the antisense direction. An antisense RNA transcribed from the antisense nucleotide sequence of the invention comprises in the molecule two or more successive repeats of an RNA sequence complementary to the part of mRNA coding for a protein of interest. Therefore, the transcribed antisense RNA will form a pair with the strand of the mRNA coding for the protein of interest, thereby enhancing the possibility of suppressing the expression of said protein in vivo. Accordingly, the antisense RNA comprising the repeats of a complementary RNA sequence is more effective in suppressing the synthesis of the protein of interest than antisense RNA having no such repeats of a sequence unit.

[0012] According to further aspects of the invention, there are provided an expression vector having the antisense nucleotide sequence of the invention, a transformant obtained by transformation with said expression vector, and a method of suppressing the intracellular expression of a protein encoded by a structural gene which includes the step of introducing the antisense nucleotide sequence of the invention into a genomic gene in a target cell.

[0013] The invention will now be described in detail.

[0014] According to a first aspect of the invention, there is provided an antisense nucleotide sequence in which two or more repeats of a structural gene of interest or a sequence representing a part of said structural gene are linked in succession in the antisense direction.

55 [0015] The term "a structural gene of interest" as used herein means a gene coding for a protein whose expression is desired to be suppressed. The protein to be under expressed is not limited to any particular type as long as it is produced in vivo; however, proteins that are expressed abundantly in vivo are preferred targets since the antisense nucleotide sequence of the invention will work more effectively. Examples of such proteins are storage proteins in plant

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seeds. More specific examples include glutelin, prolamin, globulin, albumin and so forth in cereals; particularly, rice glutelin, wheat glutenin, maize zein and barley hordein occur abundantly in seeds. Other examples include soybean conglycinin, kindney bean phaseolin, as well as potato patatin and sweet potato sporamin.

[0016] In the Examples that are given hereinafter, glutelins A and B, storage proteins in rice, were used but they are just intended as illustrative embodiments of the invention.

[0017] In the invention, the entire sequence of a particular structural gene may be employed or, alternatively, a sequence representing a part of said gene may be employed. In the latter case, a sequence from around the 5' side is preferably used because a complementary sequence corresponding to the initiation site of the structural gene is more effective in suppressing the expression of the protein of interest.

[0018] If a partial sequence is used, its size is not limited to any particular length; generally, a sequence length of at least 45 nucleotides is preferred, with a length of at least 300 nucleotides being particularly preferred (Tada et al. (1996) Breeding Science 46, 403-407).

[0019] The antisense nucleotide sequence of the invention is such that multiple units of the above-described sequence are linked in succession in the antisense direction.

[0020] The term "antisense direction" as used herein means such a direction that at least a part of the antisense DNA sequence is oriented to provide a region which is complementary to the genomic DNA of host cells. The transcript from the complementary gene has a sequence that is complementary to the RNA sequence, particularly mRNA, which is endogenous in the host.

[0021] The term "multiple units" as used herein means at least two units, preferably at least 4 units, more preferably at least 8 units.

[0022] The term "in succession" as used herein means that adjacent units of the sequence are not interrupted by any other nucleotide or that any intervening sequence is not a promoter or other sequences that will cause significant effects on the expression of the structural gene.

[0023] In the invention, multiple units of the antisense nucleotide sequence can be linked by any customary methods used in genetic engineering technology and appropriate methods may be applied without any particular limitations.

[0024] In an exemplary method, a structural gene of interest is cloned in a plasmid and the sequence containing a full length of the structural gene or a fragment thereof, containing a part of the structural gene (such sequence or a fragment thereof is referred to as an insert) is excised from a plasmid. Different restriction sites are introduced at the two terminal ends of the excised insert by a suitable means such as the addition of linkers. In this case, it is important that the sequence at the cloning site of the vector (e.g. plasmid vector) in the subsequent cloning step is taken into consideration, in order to ensure that the restriction sites at the two terminal ends of the insert may be designed in such a way that the resulting plasmid contains the structural gene of interest in the antisense direction. Subsequently, the vector having the different restriction sites at the two ends and the insert having the corresponding restriction sites at the two terminal ends are ligated to produce a gene construct containing one antisense nucleotide sequence.

[0025] In the next step, a construct is linearized by treatment with a restriction enzyme and different restriction sites are introduced at the two terminal ends. In the same manner as described in the preceding paragraph, another insert containing a full or partial length of the structural gene is excised and the corresponding restriction sites are introduced at the two terminal ends of the insert. The resulting sequence is ligated with the linearizied construct to thereby produce a gene construct containing two antisense sequences.

[0026] The same procedure may be repeated as many times as is required to produce a gene construct containing a desired number of antisense nucleotide sequences. It is also possible to start from a gene construct containing two antisense nucleotide sequences, and excise a fragment containing the antisense sequences to be used as an insert thereby to obtain a gene construct containing four antisense sequences through one ligation step. Similarly, starting from a gene construct containing four antisense sequences, a gene construct containing eight antisense sequences can be prepared through one ligation step.

[0027] In one embodiment, the invention also provides an antisense nucleotide sequence which comprises two or more different types of structural genes or sequences representing a part of them, as the structural genes of interest. Stated specifically, if different types of structural genes are to be underexpressed simultaneously, a transforming gene construct may be so designed that it comprises a linkage of multiple units of transforming gene constructs each comprising an antisense nucleotide sequence of the invention that is placed downstream of a promoter and followed by a terminator. Since each of said gene construct units comprises an antisense sequence of a different type of structural gene or a part thereof oriented in the antisense direction, multiple structural genes can be suppressed from expression by use of a single transforming gene construct.

[0028] According to a second aspect of the invention, there is provided an expression vector having the antisense nucleotide sequence of the invention.

[0029] In constructing the expression vector, regulator such as a promoter and a terminator, and a transformation vector may be appropriately selected from among those which are customarily used in genetic engineering and they are not limited to any particular types.

- [0030] Exemplary promoters include glutelin promoter, conglycinin promoter, phaseolin promoter, ADH promoter, heat shock protein promoter, a tissue specific promoter, a promoter associated with fruit ripening, prolamin promoter, RUBP carboxylase small subunit promoter cauliflower mosaic virus promoter, and so forth.
- [0031] Exemplary terminators include nopalin synthese terminator, cauliflower mosaic virus terminator, and so forth.
 [0032] Exemplary vectors include pUC plasmid vector, pBR plasmid vector, Ti plasmid vector, Ri plasmid vector, and so forth.
 - [0033] According to a third aspect of the invention, there is provided a transformant produced by transformation with the expression vector of the invention.
- [0034] The organisms to be transformed are not limited to any particular types and may be selected as appropriate from among plants, animals, microorganisms and so forth. In the present invention, transformants are preferably plants, in particular, higher plants such as monocotyledons and dicotyledons; particularly preferred are cereals (e.g. rice, wheat, maize and barley), soybean, kidney bean and potato and sweet potato.
- [0035] According to a fourth aspect of the invention, there is provided a method in which intracellular expression of proteins encoded by structural genes is suppressed by introducing the antisense nucleotide sequence of the invention into a genomic gene in a target cell.
 - [0036] The method of introducing the antisense nucleotide sequence of the invention into a genomic gene in a target cell is not limited in any particular way and a suitable transforming technique may be employed depending upon a specific object.
- [0037] For example, the antisense nucleotide sequence of the invention may be introduced into genomic genes in plant cells by the use of Agrobacterium tumefaciens, electroporation into protoplasts, liposome fusion, microinjection, and so forth.
 - [0038] Plant cells into which the antisense nucleotide sequence has been sucessfully introduced can be selected by a suitable technique such as screening of antibiotic resistant cells. The thus transformed plant cells may be cultured for regeneration into intact plants. The regenerated plant can be fixed in a variety by customary breeding techniques.
- [0039] The antisense nucleotide sequence may be introduced into genomic genes in animal cells by electroporation, liposome fusion, microinjection and other suitable methods.
 - [0040] The antisense nucleotide sequence may be introduced into genomic genes in microorganism cells by the calcium method, electroporation and other suitable methods.
- [0041] The following examples are provided for the purpose of further illustrating the present invention but are in no way to be taken as limiting.

EXAMPLES

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- [0042] In the examples, an attempt was made to suppress the storage proteins of rice, glutelin, by way of antisense RNAs in order to reduce the glutelin content in rice endosperm. With regard to glutelin, rice contains 10 or more genes in the genome coding for either glutelin A or glutelin B. The homology of their nucleotide sequences is 65%. Reducing the glutelin content of rice provides the rice with better quality for brewery.
 - [0043] A brief summary of the steps employed in the examples is as follows. Where no specific indication is made, the methods are in accordance with Maniatis T. et al. (Molecular Cloning, Cold Spring Harbor (1982)).
 - (a) The full length cDNAs of the rice storage proteins, glutelin A and glutelin B were isolated (Okita et al. (1989). Journal. Biochemical Chemistry 264: 12573-12581). The entire nucleotide sequences of the cDNAs of glutelin A and glutelin B are shown as SEQ ID NO: 1 and SEQ ID NO: 2, respectively.
 - (b) A plurality of the 312 bps sequence from the 5' upstream of the full length cDNA of glutelin A were linked in the antisense direction.
 - (c) The antisense gene comprising the linkage was used as the template for in vitro transcription to prepare the corresponding antisense RNA.
 - (d) The antisense RNA was mixed with the sense RNA of glutelin A and the mixed solution was used in in vitro translation to study the amount of glutelin A synthesis.
 - (e) The first intron of castor bean catalase gene (Ohta S. et al. (1990) Plant Cell Physiol. 31, 805-813) was introduced downstream to the glutelin promoter; and 8 repetitions of the 312 bps sequence from the 5' upstream of the full length cDNA of glutelin A were placed downstream to said intron. A terminator of nopaline synthase was further added to provide an expression plasmid vector. Similarly, an antisense gene comprising 8 repetitions of the 287 bps sequence from the 5' upstream of the full length cDNA of glutelin B was prepared, which was then introduced into said expression plasmid vector.
 - (f) The expression plasmid vector was used to transform a rice plant and the glutelin content in the rice seeds was studied.

Example 1. In vitro effects of tandem-repeated antisense RNA

- (1) RNA synthesis in vitro
- (1-1) Synthesis of sense mRNA for glutelin A

[0044] A full-length glutelin A cDNA was inserted, in the sense direction, into the multiple cloning site EcoRI/BamH1 that is located downstream of the T7 promoter in Bluescript plasmid(product from Toyobo Co.) (Fig. 1, Glu sense RNA). Transcription was carried out in a reaction mixture containing the plasmid described above (5 μg of DNA in 50 μl) by adding thereto 20 mM each of ATP, CTP, and GTP (4 μl), 2.5 mM GTP (1 μl), 5 mM Cap analogue (5 μl), 0.5 M DTT (1 μl), RNase inhibitor (50 units), and T7 RNA polymerase (10 units). The reaction mixture was incubated for 30 min at 37°C and further 4.5 hr at 37°C after addition of 20 mM GTP (1 μl).

(1-2) Synthesis of glutelin A antisense RNA

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[0045] A full-length glutelin A cDNA was inserted, in the antisense direction, to the multiple cloning site BamHVEcoRI that is located downstream of the T7 promoter in Bluescript plasmid(product from Toyobo Co.) (anti-full RNA). Similarly, a 312 bps fragment from the 5'-end of glutelin A cDNA sequence was inserted in Bluescript plasmid in the antisense direction, singly (anti x 1 RNA), 4-times repeatedly (anti x 4 RNA), or 8-times repeatedly (anti x 8 RNA) (Fig. 1) as fol-20 lows. First, a Xbal linker (CTCTAGAG) was added to the Stul site that existed at 312 bps downstream from the 5'-end of glutelin A cDNA. Plasmid anti x 1 RNA was constructed by inserting the BamHVXbal fragment thereof into the multiple cloning site Xbal/BamHI, which is located downstream of the T7 promoter, in Bluescript plasmid (product of Toyobo Co.). Next, a BamHI linker (CGGATCCG) was added to the Stul site which is located 312 bps downstream from the 5'-end of glutelin A cDNA, and the BamHVScal fragment separated therefrom was inserted into the BamHVSmall site of the above plasmid anti x 1 RNA, to provide plasmid anti x 2 RNA. The plasmid was digested at the Xbal site, treated with DNA polymerase to fill in the sticky ends, and subsequently treated with EcoRI to provide a fragment containing two of said 5'end sequence of appoximately 312 bps. Separately, plasmid anti x 2 RNA was digested at the PstI site, filled in by DNA polymerase, and cleaved with EcoRI. Plasmid anti x 4 RNA was constructed by inserting the above fragment into this EcoRI site. Similarly, plasmid anti x 8 RNA was constructed by inserting the XhoI fragment which was removed from plasmid anti x 4 RNA after the Xbal cut site was filled in by treatment with DNA polymerase into the HinclVXhol site of plasmid anti x 4 RNA.

[0046] Transcription was carried out in a reaction mixture that contained the plasmid (5 μg of DNA), a mixture of 20 mM each of ATP, CTP, UTP, and GTP (5 μl), 0.5 M DTT (1 μl), RNase inhibitor (50 units), and T7 RNA polymerase (10 units), by incubation at 37°C for 5 hr.

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(2) In vitro translation reaction by using wheat germ extract

[0047] Translation experiments were conducted by using wheat germ extract (Amersham). To the Glu sense RNA (2 picomoles) that had translation activity to glutelin was added antisense RNA against glutelin sense RNA as follows: anti-full RNA, anti x 1 RNA, anti x 4 RNA, or anti x 8 RNA, to provide an amount of 0.2, 0.5, 1, and 2 picomoles, respectively. Translation reaction was carried out at 30°C for 1 hr after addition of 15 µl of wheat germ extract to a mixture containing 2 µl of 1 mM amino acid mixture (19 amino acids lacking methionine) and 0.5 µl of 35S-methionine (1,000 Ci/mmol). The reaction was terminated by addition of 20% SDS (2.5 µl) followed by heating at 95°C for 5 min. The translation products were separated by electrophoresis on 13% polyacrylamide gel. The radioactivity was detected and measured by using Imaging analyzer (Fuji Film Co.).

(3) In vitro analysis of the effect of antisense RNA

[0048] Amount of the translation product synthesized by wheat germ extract, in the presence of the antisense RNAs, was analyzed by using Imaging analyser (Fig. 2). No band appeared if no RNA was added (lane 1), while the glutelin band appeared when the glutelin sense RNA alone was added (lane 2). However, the intensity of the glutelin band was smaller as the moles of the antisense RNA added was greater (lanes 3-6).

[0049] From the radioactivity of the glutelin bands, amounts of glutelin synthesized under the given conditions were determined. The graph (Fig. 3) shows the results for the amounts of glutelin synthesized under the given conditions, in percentage as compared with the control which contained no antisense RNA. The decrease of glutelin depended on the number of the linked 5'-upstream 312 bps sequences of the glutelin A cDNA. Such decrease was larger in cases in which the antisense RNA comprised 4-tandem repeats or 8-tandem repeats of the 5'-upstream 312 bps sequence than in the case in which the full-length glutelin antisense RNA was used.

Example 2. Production and analysis of transgenic plants transformed by glutelin A antisense gene

- (1) Construction of 8-tandem repeats of glutelin A antisense gene for transformation
- [0050] A SmallXbal fragment containing the first intron of the castor bean catalase gene (Ohta S. et al. (1990) Plant Cell Physiol. 31, 805-813) was inserted into the ScallXbal site downstream of the glutelin promoter (Takaiwa et al. (1987) FEBS Lett. 221:43-47), and then a XballSacl fragment containing 8-tandem repeats of the 5'-upstream 312 bps from the glutelin A cDNA was inserted into its downstream, in the antisense direction. A ScallEcoRl fragment containing the terminator of nopalin synthase (Depicker et al. (1982) J. Mol. Appl. Genet. 561-573) was joined to it and then the so formed antisense gene was inserted into a plasmid vector having a hygromycin resistance gene, to generate pSBHCl x 8A (Fig. 4). The intron inserted in the plasmid enhances the expression of glutelin A antisense gene.
 - (2) Construction of a full-length glutelin A antisense gene for transformation
- 15 [0051] A SmallXbal fragment containing the first intron of the castor bean catalase gene (Ohta S. et al. (1990) Plant Cell Physiol. 31, 805-813) was inserted into the ScallXbal site downstream of the glutelin promoter, and then the ScallXbal fragment containing glutelin A cDNA was inserted into its downstream, in the antisense direction. A ScallEcoRI fragment containing the terminator of nopalin synthase was joined to it and then the so formed antisense gene was inserted into a plasmid vector having a hygromycin resistance gene, to generate pSBHCI-FA (Fig. 4).
 - (3) Transformation to rice plants

[0052] A variety of the rice plant, "Tsukinohikari", was transformed by Agrobacterium tumefaciens LBA4404 which harbored the above plasmid pSBHCI x 8A or pSBHCI-FA, according to the method of Hiel et al. (Plant J. 6, 271-282 (1994)). Transformed calluses were selected in the presence of hygromycin as described by Hiel et al. (ibid).

(4) Preparation of DNA and RNA

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[0053] DNA of transgenic plants was prepared from redifferentiated seedlings before they were transferred to pot cultivation in a closed greenhouse.

[0054] RNA was prepared by the SDS-phenol method (Uchimiya et al. Manual for plant genetic engineering, Kodan-sha Co., Tokyo, Japan).

(5) PCR and Northern analyses of transformed plants

[0055] Transformation was confirmed by detecting the glutelin antisense gene after the gene was amplified by PCR. Genomic DNA (120 ng), four dNTPs (200 µM each), primers (10 pmol/reaction), and Taq DNA polymerase (Takara Biomedicals Co.) (1 unit) were mixed. The mixture was heated for 2 min at 94°C; 50°C, and 72°C, respectively, in the first cycle, and subsequently 1 min at each temperature for 34 cycles. For amplification of the 8-tandem repeats of the glutelin antisense gene, the following primers were used:

5'-AGTGGGCTGCAGGAATTCGATATCAAGCTT-3' (SEQ ID NO:3)

5'-AGTACATAGCAGCAAAACAT-3' (SEQ ID NO:4)

Similarly, for amplification of the full-length glutelin A antisense gene, the following primers were used: 5'-TACATAGCTTTAACTGATAATCTGA-3' (SEQ ID NO:5)

5'-AGTACATAGCAGCAAAACAT-3' (SEQ ID NO:6)

[0056] Northern analysis was performed according to a conventional procedure after total RNA (20 µg) was separated by electrophoresis on 1% agarose gel. The total RNA was prepared, for this analysis, from immature seeds 8 days after flowering of the transformant which was confiremed to have the 8 tandem-repeats of the glutelin A antisense gene, or those of the control plant that was transformed with the hygromycin resistance gene alone. The probe was the 5-upstream 312 bps sequence derived from glutelin A cDNA.

[0057] The probe, (25 ng of a DNA fragment) was labeled with ³²P by using Ready Prime DNA Labeling System (Amersham), before the Northern analysis.

(6) Production of transformants with 8 tandem-repeats of the glutelin A antisense gene

[0058] Individual 20 redifferentiated lines were obtained by regeneration of hygromycin-resistant calluses that had been transformed with the 8-tandem repeats of glutelin A antisense gene. Transformation by the antisense gene was confirmed by PCR followed by agarose electrophoresis as shown in Fig. 5. A band appeared, as expected, at the posi-

tion of approximately 1.2 kbps, in the lane where the plasmid containing the antisense gene was used as a template in the PCR (Fig. 5, lane 2). All 20 transformed lines gave the same band at 1.2 kbps (Fig. 5, lanes 3-22). No such band was detected when the DNA, from untransformed lines, was used for a template in the PCR reaction (Fig. 5, lane 23). [0059] Accordingly, we concluded that the above 20 transformants carried the 8-tandem repeats of the glutelin antisense gene.

(7) Production of transformants with full-length glutelin antisense gene

[0060] Individual 18 redifferentiated plant lines were obtained by regeneration from hygromycin-resistant caltuses that had been transformed with the full-length glutelin A antisense gene. Fig. 6 shows the results of agarose electrophoresis after PCR to confirm the introduction of the antisense gene. A band appeared, as expected at the position of approximately 1.7 kbps, in the lane where the plasmid containing the antisense gene was used as a template (Fig. 6, lane 2). All 18 transformants gave the same band at 1.7 kbps (Fig. 6, lanes 3-20). No such band was detected when the DNA, from untransformed lines, was used for a template in the PCR reaction (Fig. 6, lane 21).

15 [0061] Accordingly, we concluded that the above 18 transformants carried the full-length glutelin A antisense gene.

(8) Analysis of transcription product in immature seeds from transformed plants

[0062] Fig. 7 shows the results of Northern analysis for control five lines and ten transformed lines. The transformed lines carrying the 8-tandem repeats of the glutelin A antisense gene produced apparently less amount of glutelin A mRNA (Fig. 7, lanes 6-15) than the control five lines (Fig. 7, lanes 1-5). Table 1 shows the intensity of signals as determined by Bio-imaging analyzer BAS1000 (Fuji Film Co.). The results are expressed in percentage of the mean value from the five control lines. Thus, we confirmed that the content of glutelin A mRNA was significantly lower in immature seeds of the transformants, by the effect of the introduced 8-tandem repeats of the glutelin antisense gene.

Table 1

	Level of transcription product in immature seeds													
		Lane number												
Control	6	7	8	. 9	10	11	12	13	14	15				
100	31.1	11.9	13.1	11.7	10.0	7.7	8.2	3.8	3.0	1.2				

(9) Measurement of protein content in the seeds

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[0063] Unpolished seeds from self-fertilized rice plants were ground and the proteins were extracted from 50 mg of the powder. The proteins were separated by electrophoresis on 14% polyacrylamide gel, visualized by Coomassie blue staining, and measured with a densitometer Model GS-670 (Bio-Rad Laboratories) for relative glutelin content to that of control plants.

[0064] The results were expressed in percentage of the glutelin content in self-fertilized seeds of untransformed, control rice plants (Tables 2 and 3).

[0065] The glutelin content was decreased to 63.2%, on average, by the transformants with the 8-tandem repeats of the glutelin A antisense gene and to 75.4% by those with the full-length glutelin A antisense gene.

[0066] Moreover, statistical analysis confirmed that the glutelin content was significantly lower in the transformants with the 8-tandem repeats of the glutelin A antisense gene than those with the full-length glutelin A antisense gene (p < 0.10).

Table 2

Glutelin level in transformed plants (percentage of control) Transformants with 8-tandem repeats of glutelin antisense gene												
Line number Glutelin content Line number Glutelin content												
1	59.9	11	66.0									
2	66.9	12	76.2									

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Table 2 (continued)

Glutelin level in transformed plants (percentage of control) Transformants with 8-tandem repeats of glutelin antisense gene												
Line number	Glutelin content	Line number	Glutelin content									
3	42.3	13	48.1									
4	42.6	14	83.4									
5	66.6	15	71.7									
6	77.0	16	98.2									
. 7	39.7 ·	17	67.8									
8	40.1	18	61.0									
9	55.5-	19	. 93.2									
10	46.1	20	61.2									

10

15

Table 3

	ntent in transformed nants with full-length	•	•
Line number	Glutelin content	Line number	Glutelin content
1	95.2	10	77.9
2	38.9	11	59.3
3	107.2	12	60.3
4	56.3	13	41.2
5	96.2	14	96.2
6	110.6	15	60.5
. 7	73.6	16	114.3
8	72.1	17	47.4
9	66.6	18	83.0

Example 3. Production and analysis of transgenic plants with both glutelin A and glutelin B antisense genes

- (1) Construction of 8-tandem repeats of glutelin AB antisense gene for transformation
- [0067] A SmallXbal fragment, which contained the first intron of the castor bean catalase gene (Ohta S. et al. (1990) Plant Cell Physiol. 31, 805-813), was inserted into the ScallXbal site downstream of the glutelin promoter. Then the XballSphl fragment, which contained the 8-tandem repeats of the 5'-upstream 287 bps from the full-length glutelin B cDNA, was linked downstream to the above sequence, in the antisense direction. After a SphllHindIII fragment containing the terminator of nopaline synthase was linked, the sequence was inserted into the HindIII site of plasmid vector pSBHCI x 8A, constructed in Example 2, to generate pSBHCI x 8AB (Fig. 8).
 - (2) Construction of full-length glutelin AB antisense gene for transformation
- [0068] A SmallXbal fragment, which contained the first intron of the castor bean catalase gene (Ohta S. et al. (1990)
 Plant Cell Physiol. 31, 805-813), was inserted into the ScallXbal site downstream of the glutelin promoter. Then, a
 SphllXbal fragment, which contained the full-length glutelin B cDNA, was linked downstream to the above sequence,
 in the antisense direction. After a SphllHindIII fragment containing the terminator of nopaline synthase was linked, the
 sequence was inserted into the HindIII site of plasmid vector pSBHCI-FA, constructed in Example 2, to generate pSB-

HCI-FAB (Fig. 8).

(3) Transformation to rice plants

[0069] A variety of the rice plant, Tsukinohikari, was transformed by Agrobacterium tumefaciens LBA4404 which harbored the above plasmid pSBHCI x 8AB or pSBHCIFAB, according to the method of Hiel et al. (Plant J. 6, 271-282 (1994)). Transformed calluses were selected in the presence of hygromycin as described by Hiel et al.

(4) Measurement of protein content in seeds

[0070] Unpolished seeds from self-fertilized transformed rice plants were ground and the proteins were extracted from 50 mg of the powder. The proteins were separated by electrophoresis on 14% polyacrylamide gel, visualized by Coomassie blue staining, and the relative content of glutelin to that of control plants was measured with a densitometer-Model GS-670 (Bio-Rad Laboratories).

(60071) The results were expressed in percentage of the glutelin content in the self-fertilized seeds of the untransformed, control rice plants (Table 4).

[0072] The glutelin content was reduced to 57.1%, on average, by the transformants with the 8-tandem repeats of glutelin AB antisense gene and to 69.3% by those with the full-length glutelin AB antisense gene.

[0073] Moreover, statistical analysis confirmed that the glutelin content was significantly lower in the transformants with the 8-tandem repeats of glutelin AB antisense gene than those with the full-length glutelin AB antisense gene (p<0.10).

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Gitutelin content ,							
•	Mean						
Untransformed lines	100						
Transformed lines with 8-tandem repeats of glutelin AB antisense gene	57.1						
Transformed lines with the full-length glutelin AB antisense gene	69.3						

Example 4. Production and analysis of transformant with glutelin A antisense gene

(1) Construction of 2-tandem repeats and 4-tandem repeats of glutelin A antisense gene for transformation

[0074] A SmallXbal fragment, which contained the first intron of the castor bean catalase gene (Ohta S. et al. (1990). Plant Cell Physiol. 31, 805-813), was inserted into the ScallXbal site downstream of the glutelin promoter. Then, a XballScal fragment, which contained 2-tandem repeats or 4-tandem repeats of the 5'-upstream 312 bps from the full-length glutelin A cDNA, was linked downstream to the above sequence, in the antisense direction. After a ScallEcoRI fragment containing the terminator of nopaline synthase was linked, the sequence was inserted into the transformation plasmid vector with the hygromycin resistance gene, thereby to generate pSBHCI x 2A and pSBHCI x 4A, respectively.

(2) Transformation to rice plants

[0075] A variety of the rice plant, Tukinohikari, was transformed by Agrobacterium tumefaciens LBA4404 which harbored the above plasmid pSBHCl x 2A or pSBHCl x 4A, according to the method of Hiel et al. (Plant J. 6, 271-282 (1994)). Transformed calluses were selected in the presence of hygromycin as described by Hiel et al. (ibid).

(3) Measurement of protein content in seeds

[0076] Unpolished seeds of self-fertilized transgenic rice plants were ground and the proteins were extracted from 50 mg of the powder. The proteins were separated by electrophoresis on 14% polyacrylamide gel visualized by Coornassie blue staining, and measured with a densitometer Model GS-670 (Bio-Rad) for the relative content of glutelin to that of control plants. The result was expressed in percentage of the glutelin level in the self-fertilized seeds of the untransformed, control rice plants (Tables 5 and 6).

[0077] The glutelin content was decreased to 71.7%, on average, by the transformants with the 2-tandem repeats of

the glutelin A antisense gene and to 71.5% by those with the 4-tandem repeats of the glutelin A gene.

Table 5

Glutelin content in transgenic plants (percentage of control) Transformants with 2-tandem repeats of the glutelin A antisense gene										
Line number	Glutelin content									
1	66.5									
2	66.2									
3	. 67.5									
4	93.8									
5	64.2									
6	60.6									
7	75.0									
8	79.5									
Mean	71.7									

25

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15

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Table 6

68.0

66.7

67.0

73.1

67.0

71.5

•	Glutelin content in transgenic plants (percentage of control) Transformants with 4-tandem repeats of the glutelin A antisense gene						
	Line number	Glutelin content					
25	1	54.7					
	2	89.0					
	3	93.9					
	4	64.1					
· ·	5	68.0					

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EFFECT OF THE INVENTION

[0078] The antisense nucleotide sequence in the present invention represents a single copy of tandem repeats of the structural gene of interest or a fragment thereof which are successively linked in the antisense direction. The antisense 85 RNA transcribed from the antisense nucleotide sequence consists of plural repeats of the RNA sequence each being complementary to the corresponding part of the mRNA of the target protein. Therefore, the antisense RNA has a higher ability to hybridize with the mRNA and to suppress the in vivo expression of the protein. Thus, the antisense RNA of the invention containing such tandem repeats is superior to an antisense RNA without any such repeats in suppressing the

8

Mean

synthesis of target proteins.

[0079] The description has been made mainly on the basis of suppression to the synthesis of plant proteins. Needless to say, however, the antisense sequence of the present invention is useful for suppressing the synthesis of animal proteins such as specific milk proteins or specific proteins for the purpose of treating immunological or pathogenic diseases.

SEQUENCE LISTING

					500												
5	(2)	(i)	SEQ A) B) O)	UENC L) LE L) TY L) ST	E CE NGTH PE: RAND	ARAC	TERI 44 b elc SS:	STIC ase acid doub	S: pair	· s ·							
10		•						A to	mRN	A						•	•
		(vi)		GINA ORG		ORCE 4:	:										•
15		(ix)	FEA	TURE	: ·G1	utel	in A	CDN	A								
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20	>=>	~~~		mm<	a cia	~**	* CC	#				***				~ ₹3	102
												TGC Cys					103
25										_	_	TGG					151
·	Ala	Gln 25	Gln	Leu	Leu	Gly	GLn 30	Ser	Thr	Ser	Gln	Trp 35	Gln	Ser	Ser	Arg	•
												TTG				_	199
30	Arg 40	Gly	Ser	Pro	Arg	GLu 45	Cys	YEG	Phe .	Asp	Arg 50	Leu	Gln	Ala	Phe	55	
					_					•		ACT					247
	SLO	Ile	Arg	Ser	Val 60	Arg	Ser	Gin	Ala	GLY 65	Thr	Thr	Glu	Phe	Phe 70	Asp	
35 .	CTC.	тС-т	BBT	GNG	CBB	-	CAA	ጥርሞ	»cc	CCA	CT.	TCT	·CTT	CTC.	CCT	CCA	295
												Ser					233
	GTT	ATT	GAA	CCT	AGA	GGC	CTT	CTA	CTA	CCC	CAT	TAC	ACT	AAT	GGT	GCA	343
40	Val	Ile	G1u 90	Pro	Arg	Gly	Leu	Leu 95	Leu	Pro	His	Tyr	Thr 100		Gly	Ala	
												ACA					391
45	Ser	Leu 105		TYT	He	Ile	G1a 110	Gly	Arg	GLY	Ile	Thr 115	Gly	Pro	Thr	Phe	•
	CCA	ccc	TCT	CCT	GAG	TCC	TAC	CBA	CAA	CNG	ጥተር	CAA	·CBB	TCA	GGC	CAA	439
		Gly										Gln				Gln 135	
<i>50</i>	GCC	CAA	TTG	ACC	GAA	AGT	CAA	AGC	CAA	AGT	CAA	AAG	TTC	AAG	GAT	GAA	487
											Gln					Glu	

E				_					CAA Gln 160	•			Ile	_			535
5	_								AAT Asn								583
10									AAC Asn			_					, 631
15									GGA Gly								679
						Glu			TCA Ser							_	727
20									CTT Leu 240						_		775
25				_		-			CAA Gln								823
30									Pro				Leu				871 .
				_		_									_	TAT Tyr 295	919
35 ,						Gly			TGC	-	Asn		-			Thr	967
40					Arg					Ile					Arg	GCT Ala	1015
				Asn	_				Arg				-	Ast		CAG Gln	1063
45			Pro					Va]			-		Val			TAA J neA J	1111
<i>50</i>		Tyr					Leu					Ast				CAC A His 375	1159
	AGC	GTC	GIG	TAT	ATT	ACT	CÃ	GGC	CG1	GCC	CGG	GT:	CA	A GT	c Gt	CAAC	1207

•

	Ser	Val	Val	Tyr	11a 380	Thr	Gln	Gly	Arg	Ala 385	Arg	Val	Gln	Val	Val 390	Asn	
5						GTG Val											1255
10						CAC								Gln			1303
15						GCA Ala										AGC Ser	1351
10						AGT Ser 445										GTT Val 455	1399 .
20	_	_		_			_				Glu					AAG . Lys .	1447
25			Arg		Asp	GAG Glu										AAG Lys	1495
		Tyr				TAT Tyr								p	GGCT	TGC GĢ	1546
30	ATA	NAGA	ATA I	CŢN	VATA	AA TI	AAAT.	IGCA	R GC	Aatt	GTTT	TGĊ	T GCT	ATG	TACT	GTCCAG	1606
	TCT	rtcg	ACT I	AATG	itga:	ia ai	AGCC	TCTC	T TT	ATCC	TT		•			•	1644
35	(2)) SE(() () ()	QUENCA) LI B) T C) S	ce ci engti rpe: trani	SEQ HARAC H: 10 DEDNI DEDNI OGY:	CTER 634 leic ESS:	ISTI base aci dou	C5: pai d	rs			٠				
40			OR:		AL S	TYPE		NA t	o mR	N/A							
4 5) FE	azuri	E: G	n. lute: ESCR				ID N	o: 2	! :				-	
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50			_					Gły					Lei			r ccc	99
													•				•

5										CAA Gln							147
•				_			_			CCA Pro 55				_			195
10										GAG Glu							243
15										Val Val							291
										GGC							339
20					Met					CCC							387
25										Gln 135			-				435
										CAT His					Gly		483
30 .			_						Ala							GLY	531
35								Tyr					Asn			GCC	579
40		Gln					Gln					Leu				AAC Asn 205	627
•						Gln					Ser					CAC His	675
45					Ile					Gly	•				se:	GAG Glu	723
50				Ile					λla					Sei		TAK A	771
	GAT	CAA	AGA	GGA	GAG	ATC	ATA	CAT	GTG	, AAC	AA1	r GG(CTI	CA	A II	G TTG	819

16

	Asp	Gln 255	Arg	Gly	Glu	Ile	11e 260	His	Val	Lys	Asn	Gly 265	Leu	Gln	Leu	Leu	
5 .							CAG Gln										867
10							AGT Ser				Gln					TGG ' Trp	915
15					Glu		TTT Phe										963
· ·	GAA Glu	AAT Asn	CCT Pro 320	AGT Ser	CGT Arg	GCT Ala	gat Asp	TCA Ser 325	TAC Tyr	AAC Asn	CCA Pro	CGT Arg	GCC Ala 330	GGA Gly	AGG Arg	ATA Ile	1011
20	Thr		Val				AAG Lys 340	Phe									1059
25	AGC Ser 350	GCT	AGC Thr	AGA Arg	GTA Val	AAC Àsn 355	CTA Leu	TAC Tyr	CAG Gln	AAT Asn	GCT Ala 360	Ile	CTC	TCG Ser	CCG Pro	TTC Phe 365	1107
							AGT Ser										1155
30 •							AAC Asn										1203
35							TTG Leu										1251
			Ala										Ile			AAC Asn	1299
40		Asn										Asn				CGT Arg 445	1347
45	_										Tyr					GAG	1395
50	_									Gly					Ala	TTC	1443
																TCC	1491

		480		485			190		
5			Ser Glu		aaigtaa	TTGAGAAC	CTA GTATOGO	SCGT AGAG	1546
	TANAATAA	AA CACCI	CAAGT AT	GACACTT	G GTGGT	SATTC TG	ITCGATAT C	AGTACTAAA	1606
10	TAAAGGTT	AC AAACT	TCTIA AT	TTTCCT					1634
15		SEQUENC (A) LI (B) TY (C) SY	FOR SEQ CE CHARAC ENGTH: 30 CPE: DUCL TRANDEDNE OPOLOGY:	TERISTI base p eic aci SS: sin	CS: airs d	-	-	•	
	(ii)	MOLECUI	LAR TYPE:	Other	nucleic	acids,	synthetic	DNA.	
20			ce descri Attoga ta		_	%O: 3: 30		•	
25		SEQUENC (A) LI (B) T (C) S:	FOR SEQ CE CHARAC ENGTH: 20 YPE: nucl TRANDEDNE OPOLOGY:	TERISTI base peic aci SS: sin	cs: pairs d		•		
	(ii)	HOLECA	Lar type:	Other	nucleic	acids,	synthetic	DNA	
30	(xi) AGTACATA	_	CE DESCRI RAACAI	PTION:	SEQ ID	NO: 4: 20		•	•
35		SEQUENCE (A) LI (B) T (C) S	FOR SEQ CE CHARAC ENGTH: 25 YPE: nucl TRANDEDNE OPOLOGY:	TERISTI base p eic aci SS: sir	CS: pairs ld				•
40	(ii)	HOLECU	Lar type:	Other	nucleic	acids,	synthetic	DNA	
			CE DESCRI TGATAA TO		SEQ ID	NO: 5: 25			
4 5		SEQUEN (A) L (B) T (C) S	FOR SEQ CE CHARAC ENGTH: 20 YPE: nucl TRANDEDNE OPOLOGY:	TERIST: base page action actions action actions action	ICS: pairs id ngle				
50	(ii) HOLECU	LAR TYPE:	Other	nucleic	acids,	synthetic	DNA	
	(*1	SECUEN	CE DESCRI	י ארדראי	SEO ID	NO. S.			

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Claims

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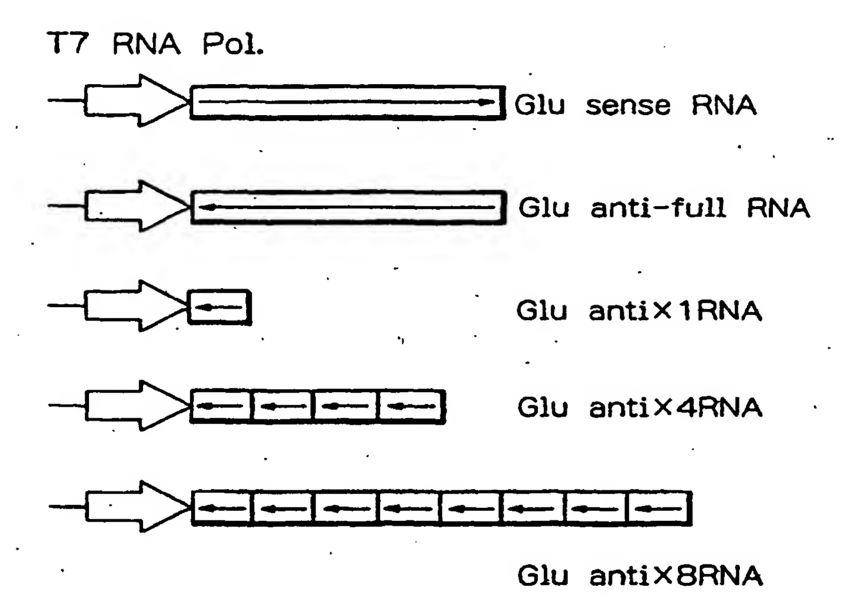
- 1. An antisense nucleotide sequence comprising two or more successive repeats of the tructural gene of interest or a fragment thereof in the antisense direction.
- An antisense nucleotide sequence as claimed in Claim 1, which comprises two or more successive repeats of the sequence of a 5' region of the desired gene of interest.
 - An antisense nucleotide sequence as claimed in Claim 2, wherein the sequence of 5' region comprises at least 45
 nucleotides.
- 4. An antisense nucleotide sequence as claimed in Claim 2, wherein the sequence of 5' region comprises at least 300 in nucleotides.
 - 5. An antisense nucleotide sequence as claimed in any of Claims 1 to 4, which comprises at least 4 repetitions of the structural gene of interest or a fragment thereof.

6. An antisense nucleotide sequence as claimed in any of Claims 1 to 4, which comprises at least 8 repetitions of the structural gene of interest or a fragment thereof.

- 7. An antisense nucleotide sequence as claimed in any of Claims 1 to 6, which comprises different sets of two or more successive repeats of the tructural gene of interest or a fragment thereof in the antisense direction.
 - 8. An antisense nucleotide sequence as claimed in any of Claims 1 to 7, wherein said gene of interest is a gene coding for a storage protein of a plant seed.
- 95. An antisens nucleotide sequence as claimed in Claim 8, wherein the storage protein is glutelin A or gluterin B.
 - 10. Án expression vector comprising an antisense nucleotide sequence of any of Claims 1-9.
 - 11. A transgenic host which has been transfromed by an expression vector of Claim 10.
 - 12. A transgenic host of Claim 11, which is a plant.
 - 13. A transgenic host of Claim 11 or 12, which is a rice plant.
- 45 14. A method for suppressing the in vivo expression of the protein of interest encoded by a structural gene comprising introducing an antisense nucleotide sequence of any of Claims 1-9 against said structural gene into the genomic gene of a target cell.

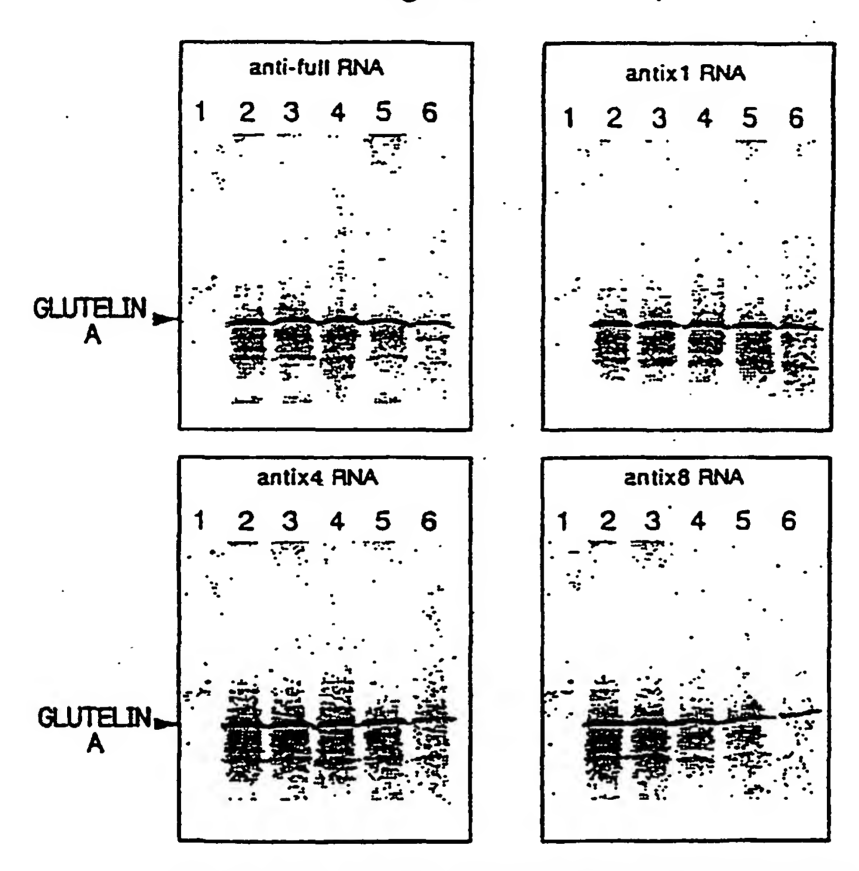
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Fig. 1



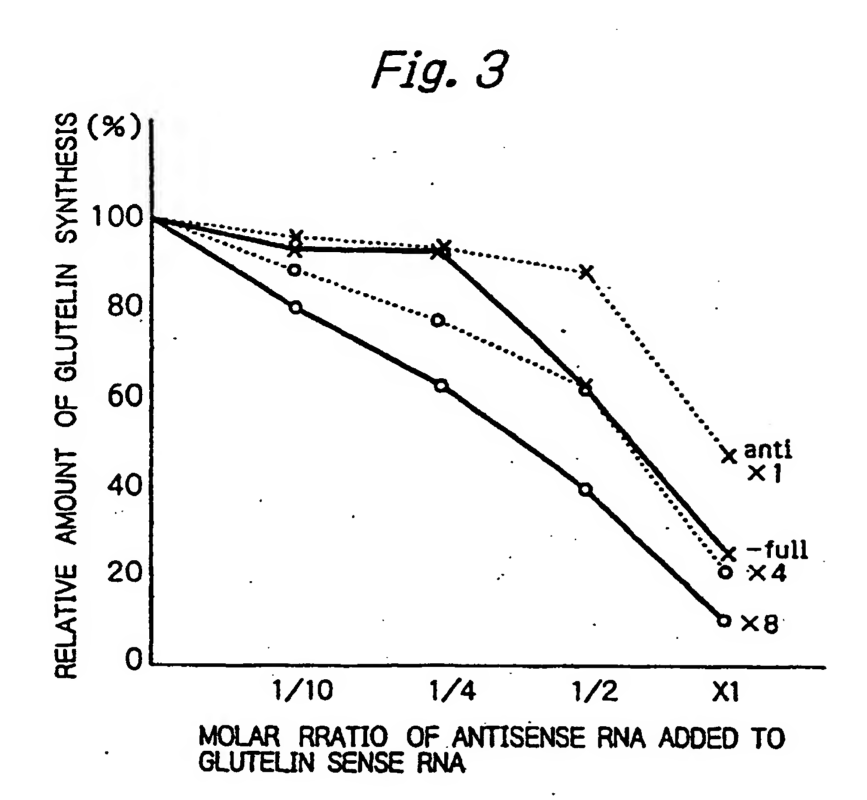
STRUCTURE OF GENES USED IN IN VITRO RNA SYNTHESYS

Fig. 2



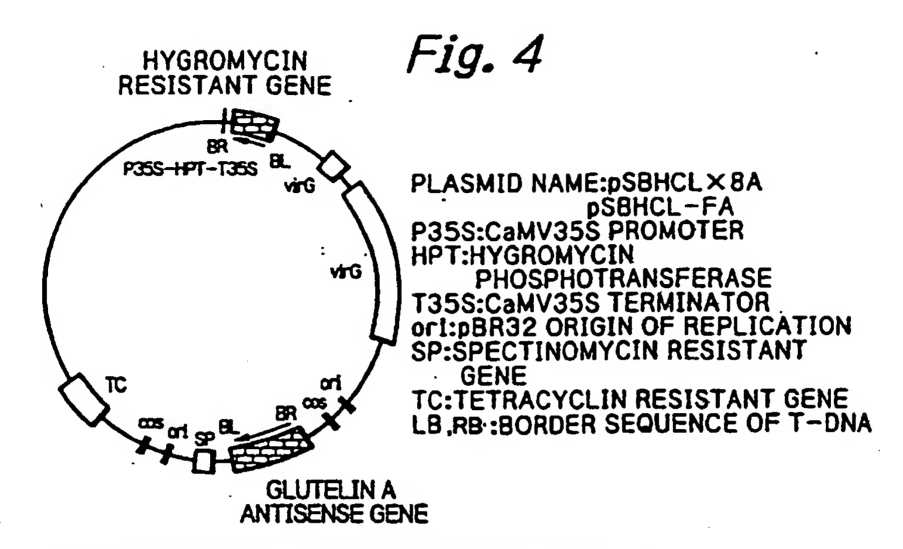
ANALYSIS OF TRANSLATION PRODUCTS IN WHEAT GERM **EXTRACT**

LANE 1 RNA NOT ADDED
LANE 2 GULUTELIN SENSE RNA ALONE (2p MOLES)
LANE 3 ANTISENSE RNA 0.2p MOLES ADDED
LANE 4 ANTISENSE RNA 0.5p MOLES ADDED
LANE 5 ANTISENSE RNA 1p MOLES ADDED
LANE 6 ANTISENSE RNA 2p MOLES ADDED

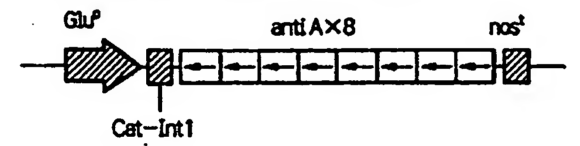


CHANGES IN GLUTELIN SYNTHESIS

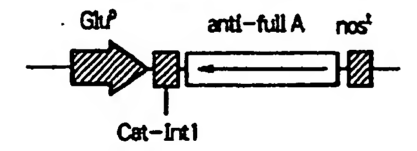
××	Glu anti×1 RNA
××	Glu anti-full RNA
oo	Glu anti×4 RNA
00	Glu anti X 8 RNA



(1) 8-REPEATS OF GLUTELIN ANTISENSE GENE(3.7kbp)



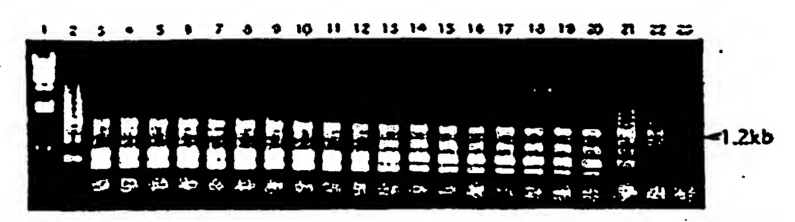
(2) FULL LENGTH GLUTELIN ANTISENSE GENE(2.9kbp)



Glu^P: GLUTELIN PROMOTER
Cat-Int1: FIRST INTRON OF CASTOR BEAN CATALASE GENE
nos¹: TERMINATOR OF NOPALIN SYNTHASE GENE
anti × 8A: 8-REPEATS OF GLUTELIN A ANTISENSE SEQUENCE
anti × fulla: FULL LENGTH ANTISENSE SEQUENCE OF GLUTRLIN A

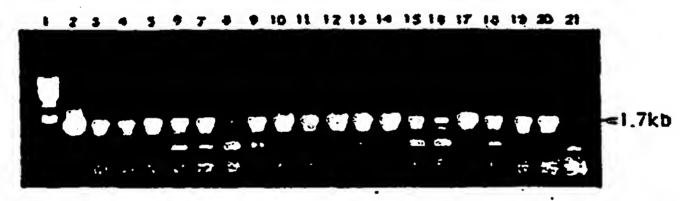
STRUCTURE OF PLASMID VECTOR USED IN TRANSFORMATION

Fig. 5



PCR ANALYSIS OF TRANSFORMANTS HAVING 8-REPEATS OF GLUTELIN A ANTISENSE GENE

Fig. 6



PCR ANALYSIS OF TRANSFORMANTS HAVING FULL LENGTH GLUTELIN A ANTISENSE GENE

Fig. 7



NORTHERN BLOT ANALYSIS OF TRANSFORMANTS HAVING 8-REPEATS OF GLUTELIN A ANTISENSE GENE

HYGROMYCIN RESISTANT GENE 98 81 P355-HPT-T355 WG

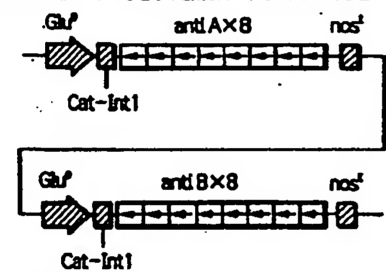
Fig. 8

PLASMID NAME:pSBHCL×8AB
pSBHCL-FAB
P35S:CaMV35S PROMOTER
HPT:HYGROMYCIN
PHOSPHOTRANSFERASE
T35S:CaMV35S TERMINATOR
ori:pBR32 ORIGIN OF REPLICATION
SP:SPECTINOMYCIN RESISTANT
GENE

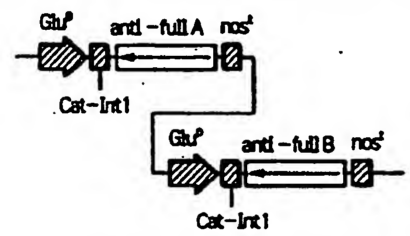
TC:TETRACYCLIN RESISTANT GENE LB,RB:BORDER SEQUENCE OF T-DNA

GLUTELIN B GLUTELIN A ANTISENSE GENE ANTISENSE GENE

(1) 8-REPEATS OF GLUTELIN A.B ANTISENSE GENE(7.4kbp)



(2) FULL LENGTH GLUTELIN A.B ANTISENSE GENE(5.8 kbp)



Glu^P: GLUTELIN PROMOTER
Cat-Int1: FIRST INTRON OF CASTOR BEAN CATALASE GENE
nos^t: TERMINATOR OF NOPALIN SYNTHASE GENE
anti×8A: 8-REPEATS OF GLUTELIN A ANTISENSE SEQUENCE
anti×8B: 8-REPEATS OF GLUTELIN B ANTISENSE SEQUENCE
anti×fulla: FULL LENGTH GLUTELIN B ANTISENSE SEQUENCE
anti×fullB: FULL LENGTH GLUTELIN B ANTISENSE SEQUENCE

STRUCTURE OF PLASMID VECTOR USED IN TRANSFORMATION

		>									
	INTERNATIONAL SEARCH REPO	RT International application No.									
		PCT/JP98/00955									
	SIFICATION OF SUBJECT MATTER										
Int.	.Cl* C12N15/29, C12N15/63, C12	N5/10, A01H1	/00, A01H5	/00							
According t	to International Patent Classification (IPC) or to both a	ational classification a	nd IPC								
B. FIELD	S SEARCHED										
Minimum d	Minimum documentation searched (classification system followed by classification symbols)										
Int.	.Clf C12N15/29, C12N15/63, C12	RS/10, A01R1.	/00, A0185	/00							
Dammarta	tion searched other than minimum documentation to th			1: 4: 6:14:							
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Electronic d	lata base consulted during the international search (nat	ne of data base and, w	here practicable, se	earch terms used)							
BIOS	SIS (DIALOG), WPI (DIALOG)		:								
				_							
C 222	MENTE CONCIDENTS DE DE DES CIANOS										
L 1000	MENTS CONSIDERED TO BE RELEVANT										
Category*	Citation of document, with indication, where ap	propriate, of the releva	of passages	Relevant to claim No.							
x	JP, 60-232092, A (The Resear	cu tonudation	or State								
A	University of New York), November 18, 1985 (18. 11. 8	E 1		1,2,5-7,10,							
Y	£ EP, 140308, A1 £ JP, 9-1		11,14								
•	4 M, 140300, RI & OF, 3-1.	32000, V		3,4,8,9,12,							
	•			1.3							
¥	TADA, Y. et al., "Environment	al risk eval	uation of	3,4,8,9,							
_	transgenic rice expressing a			12,13							
	16kDa albumin(I)", Breeding			,							
	Vol. 46, No. 4, pages 403-40		·′								
	to to the try pages to to										
Y	OKITA, T.W. et al., "Structure	and express	on of the	8,9,12,13							
	rice glutelin multigene fami:	_									
	The Journal of Biological Che	s),	į								
	Vol. 264, No. 21, pages 1257										
000			·								
Y	HIEI, Y. et al., "Efficient	transformatio	n of	12,13							
,	rice",										
	Plant Journal (1994) Vol. 6,	No. 2	j								
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T tenne	r documents are listed in the continuation of Box C.	See patent fami	ly annex.								
	categories of cited documents:			stious! Sling date or priority							
	nd defining the general state of the art which is not not to be of particular rejevence.	date and not in cor	ellict with the applicat	ion but cled to understand							
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